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APPLICATION NO.	FILING DATE	FIRST NAMED INVENTOR	ATTORNEY DOCKET NO.	CONFIRMATION NO.
10/054,710	01/22/2002	Koichi Masuda	047940-0119	5419
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MADISON, W	MADISON, WI 53701-1497		ART UNIT	PAPER NUMBER
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Please find below and/or attached an Office communication concerning this application or proceeding.

,	Application No.	Licant(s)			
	10/054,710	MASUDA ET AL.			
Office Action Summary	Examiner	Art Unit			
	Ruth A. Davis	1651			
The MAILING DATE of this communication appears on the cover sheet with the correspondence address Period for R ply					
A SHORTENED STATUTORY PERIOD FOR REPLY IS SET TO EXPIRE 3 MONTH(S) FROM THE MAILING DATE OF THIS COMMUNICATION. - Extensions of time may be available under the provisions of 37 CFR 1.136(a). In no event, however, may a reply be timely filed after SIX (6) MONTHS from the mailing date of this communication. - If the period for reply specified above is less than thirty (30) days, a reply within the statutory minimum of thirty (30) days will be considered timely. - If NO period for reply is specified above, the maximum statutory period will apply and will expire SIX (6) MONTHS from the mailing date of this communication. - Failure to reply within the set or extended period for reply will, by statute, cause the application to become ABANDONED (35 U.S.C. § 133). - Any reply received by the Office later than three months after the mailing date of this communication, even if timely filed, may reduce any earned patent term adjustment. See 37 CFR 1.704(b). Status					
1) Responsive to communication(s) filed on 22.	January 2003 .				
<u></u>	nis action is non-final.				
3) Since this application is in condition for allow					
closed in accordance with the practice under <i>Ex parte Quayle</i> , 1935 C.D. 11, 453 O.G. 213. Disposition of Claims					
4)⊠ Claim(s) <u>1-35</u> is/are pending in the application.					
4a) Of the above claim(s) 15,16,31 and 32 is/are withdrawn from consideration.					
5) Claim(s) is/are allowed.					
6)⊠ Claim(s) <u>1-14,17-30 and 33-35</u> is/are rejected.					
7) Claim(s) is/are objected to.					
8) Claim(s) are subject to restriction and/or election requirement.					
Application Papers					
9) The specification is objected to by the Examiner.10) The drawing(s) filed on is/are: a) accepted or b) objected to by the Examiner.					
Applicant may not request that any objection to the drawing(s) be held in abeyance. See 37 CFR 1.85(a).					
11) The proposed drawing correction filed on is: a) approved b) disapproved by the Examiner.					
If approved, corrected drawings are required in reply to this Office action.					
12) The oath or declaration is objected to by the Examiner.					
Priority under 35 U.S.C. §§ 119 and 120					
13) Acknowledgment is made of a claim for foreign priority under 35 U.S.C. § 119(a)-(d) or (f).					
a) ☐ All b) ☐ Some * c) ☐ None of:					
1. Certified copies of the priority documents have been received.					
2. Certified copies of the priority documents have been received in Application No					
 3. Copies of the certified copies of the priority documents have been received in this National Stage application from the International Bureau (PCT Rule 17.2(a)). * See the attached detailed Office action for a list of the certified copies not received. 					
14) Acknowledgment is made of a claim for domestic priority under 35 U.S.C. § 119(e) (to a provisional application).					
a) The translation of the foreign language provisional application has been received.					
15) Acknowledgment is made of a claim for domestic priority under 35 U.S.C. §§ 120 and/or 121.					
Attachment(s) 1) Notice of References Cited (PTO-892) 4) Interview Summary (PTO-413) Paper No(s)					
 Notice of References Cited (PTO-892) Notice of Draftsperson's Patent Drawing Review (PTO-948) Information Disclosure Statement(s) (PTO-1449) Paper No(s) 3 	5) Notice of Informal I	Patent Application (PTO-152)			

Art Unit: 1651

DETAILED ACTION

Election/Restrictions

1. Applicant's election with traverse of Group I, claims 1 – 14, 17 – 30 and 33 – 35 in Paper No. 7 is acknowledged. The traversal is on the grounds that there is not a serious burden on examiner since the groups are classified in the same class and the inventions are related. This is not found persuasive because as indicated in the previous office action, the inventions are separated and distinct as based on their separate classification (to include class and subclass). Further, even though the inventions are related, the search for one group would not necessarily anticipate or even make obvious another. It is reiterated that an overlapping search is not a coextensive search.

The requirement is still deemed proper and is therefore made FINAL.

Claims 15 - 16 and 31 - 32 are withdrawn as being drawn to non elected subject matter. Claims 1 - 4, 17 - 30 and 33 - 35 have been considered on the merits.

Claim Rejections - 35 USC § 112

- 2. The following is a quotation of the second paragraph of 35 U.S.C. 112:
 - The specification shall conclude with one or more claims particularly pointing out and distinctly claiming the subject matter which the applicant regards as his invention.
- 3. Claims 1-14, 17-30 and 33-35 are rejected under 35 U.S.C. 112, second paragraph, as being indefinite for failing to particularly point out and distinctly claim the subject matter which applicant regards as the invention.

Art Unit: 1651

Claims 1, 17 and their dependents are drawn to a method for determining effects of a test agent, however are rendered vague and indefinite because it is unclear what "effects" are being determined. Applicant fails to specifically point out what effects the method is determining, as there is no recitation of a determining step. For example, there is no defined limitation wherein a particular result indicates an "effect" or not. Moreover, it is unclear what effect, if any, must occur to practice the claimed invention.

Claims 1, 17 and their dependents are confusing for reciting both "tissue engineered cartilage matrix" and "engineered cartilage tissue" because it is unclear if applicant is using the term interchangeably. Applicant may prefer to recite the same term for clarity and consistency.

Claims 1, 17 and their dependents are rendered indefinite because while the method is drawn to determining effects on an engineered cartilage tissue (ECT), step (B) allows for the determining step to occur on isolated chondrogenic cells before they have formed into the ECT. It is unclear how one could determine the effects of an agent on an ECT by contacting cells that have yet to form an ECT.

Claims 11 and 27 are rendered vague and indefinite because it is unclear if step (C) rather comprises enzymatically degrading the ECT, or further comprises enzymatically degrading ECT.

Claims 14 and 33 are indefinite because it is unclear how one "identifies" a desirable test agent, as there is no specific property required to make it desirable.

Art Unit: 1651

Claim Rejections - 35 USC § 103

- 4. The following is a quotation of 35 U.S.C. 103(a) which forms the basis for all obviousness rejections set forth in this Office action:
 - (a) A patent may not be obtained though the invention is not identically disclosed or described as set forth in section 102 of this title, if the differences between the subject matter sought to be patented and the prior art are such that the subject matter as a whole would have been obvious at the time the invention was made to a person having ordinary skill in the art to which said subject matter pertains. Patentability shall not be negatived by the manner in which the invention was made.
- 5. This application currently names joint inventors. In considering patentability of the claims under 35 U.S.C. 103(a), the examiner presumes that the subject matter of the various claims was commonly owned at the time any inventions covered therein were made absent any evidence to the contrary. Applicant is advised of the obligation under 37 CFR 1.56 to point out the inventor and invention dates of each claim that was not commonly owned at the time a later invention was made in order for the examiner to consider the applicability of 35 U.S.C. 103(c) and potential 35 U.S.C. 102(e), (f) or (g) prior art under 35 U.S.C. 103(a).
- 6. Claims 1 8, 10, 14, 17 24, 26, 29 30 and 33 are rejected under 35 U.S.C. 103(a) as being unpatentable over Kai et al. (JP 2001 089390 A) in view of Masuda et al. (US 6197061 B1).

Applicant claims a method for determining the effect of a test agent on a tissue engineered cartilage matrix, the method comprising culturing an engineered cartilage tissue (ECT), contacting test agents with cells or tissues of the ECT, and measuring the effect that the agents have on the ECT or cells thereof. The ECT is cultured by culturing isolated chondrogenic cells for a time sufficient to form a chondrogenic cell-associated matrix, and culturing the cells with the cell-associated matrix on a semipermeable membrane in the presence of a growth factor

Art Unit: 1651

for a time sufficient to form an ECT. The cells or tissues are selected from isolated chondrogenic cells, the chondrogenic cells during culture, the cells and cell-associated matrix before or after culturing, and the ECT. Alternatively, the cells are selected from isolated chondrogenic cells, the chondrogenic cells during culture, the cells and cell-associated matrix before or after culturing, and the ECT in the presence of a known modulator of cartilage tissue. The chondrogenic cell-associated matrix comprises aggrecan, collagen types II, IX, XI, matrix proteins, and hyaluronan and the ECT comprises collagens II, IX, XI, hyaluronan, and at least about 5 mg/cc3 aggrecan, the ratio of aggrecan: hyaluronan is about 10:1 – 200:1, and the ration of aggrecan: collagen is about 1:1-10:1. The isolated chondrogenic cells are isolated from articular cartilage, costal, nasal, auricular, tracheal, epiglottic, thyroid, arytenoids or cricoid cartilages, specifically fibrocartilage from ligament, tendon, meniscus, or intervertebral disc. The chondrogenic cells are cultured on an alginate medium, the measuring is performed without addition of extrinsic radioactivity and the method further comprises identifying one or more test agents that have desirable characteristics and producing the agents as a therapeutic drug. The modulator of the ECT tissue is a matrix stimulating agent, cytokine or TNF-alpha wherein the cytokine is interleukin 1 (IL-1).

Kai teaches a method for determining effects of agents on cartilage, wherein the cartilage is cultured with IL-1 or TNF, is contacted with the test agent, and is measured for effects of the test agent (abstract). The method is used to screen for therapeutic agents (abstract).

Kai does not teach the method wherein the cartilage is cultured in the manner claimed.

However, Masuda teaches methods for culturing an engineered cartilage tissue, wherein isolated chondrocytes are cultured in alginate for a time to allow a cell associated matrix to form (col.4)

Art Unit: 1651

line 30-35). The cell associated matrix has at least about 5 mg/cc3 aggrecan, a ratio of aggrecan to hyaluronan of 10:1 to 200:1, and a ratio of aggrecan to collagen of about 1:1 to 10: (col.4 line 35-40). The chondrogenic cells are cultured on a semi-permeable membrane in the presence of growth factors (col. 4 line 44-46). The chondrocytes may be isolated from articular cartilage, fibrocartilage, costal, nasal, auricular, tracheal, epiglottic, thyroid, arytenoid or cricoid cartilages (col.5 line 31-38) and the resulting matrix comprises aggrecan, collagn II, IX, XI and hyaluronan (col.6 line 61-64). At the time of the claimed invention, it would have been obvious to one of ordinary skill in the art to culture the cartilage of Kai using the methods of Masuda because it was a known method for culturing cartilage tissues. Moreover, at the time of the claimed invention, one of ordinary skill in the art would have been motivated by common culture practices in the art to culture the cartilage of Kai via the methods of Masuda with a reasonable expectation for successfully determining the effects of test agents on cartilage cultures. Although Masuda does not teach the fibrocartilage is from a ligament, tendon, meniscus or intervertebral discs, they were each well known sources of fibrocartilage. Moreover at the time of the claimed invention, it would have been well within the purview of one of ordinary skill in the art to use any of the above sources as the source of fibrocartilage in the methods of Masuda.

7. Claims 1 - 10, 14, 17 - 26, 33 and 35 are rejected under 35 U.S.C. 103(a) as being unpatentable over Purchio et al (US 5902741) in view of Masuda.

Applicant claims a method for determining the effect of a test agent on a tissue engineered cartilage matrix, the method comprising culturing an engineered cartilage tissue (ECT), contacting test agents with cells or tissues of the ECT, and measuring the effect that the

Art Unit: 1651

agents have on the ECT or cells thereof. The ECT is cultured by culturing isolated chondrogenic cells for a time sufficient to form a chondrogenic cell-associated matrix, and culturing the cells with the cell-associated matrix on a semipermeable membrane in the presence of a growth factor for a time sufficient to form an ECT. The cells or tissues are selected from isolated chondrogenic cells, the chondrogenic cells during culture, the cells and cell-associated matrix before or after culturing, and the ECT. Alternatively, the cells are selected from isolated chondrogenic cells, the chondrogenic cells during culture, the cells and cell-associated matrix before or after culturing, and the ECT in the presence of a known modulator of cartilage tissue. The chondrogenic cell-associated matrix comprises aggrecan, collagen types II, IX, XI, matrix proteins, and hyaluronan and the ECT comprises collagens II, IX, XI, hyaluronan, and at least about 5 mg/cc3 aggrecan, the ratio of aggrecan: hyaluronan is about 10:1-200:1, and the ration of aggrecan: collagen is about 1:1-10:1. The isolated chondrogenic cells are isolated from articular cartilage, costal, nasal, auricular, tracheal, epiglottic, thyroid, arytenoids or cricoid cartilages, specifically fibrocartilage from ligament, tendon, meniscus, or intervertebral disc. The chondrogenic cells are cultured on an alginate medium, the measuring comprises measuring the amount of proteoglycan in the ECT, is performed without addition of extrinsic radioactivity and the method further comprises identifying one or more test agents that have desirable characteristics and producing the agents as a therapeutic drug. The culturing of ECT and the contacting the cells with the test agent occurs in the same well of a multi-well plate.

Purchio teaches methods for screening effects of test agents on cartilage cultures wherein the cultures are exposed to the test agents and the effects are measured (col.16 line 12-22).

Examples of such effects include the amount of proteoglycan (col.16 line 27-34). Specifically,

Art Unit: 1651

chondrocytes are harvested from articular cartilage and cultured in multi-well plates (col.21 lines 18-60) and the test agents are identified for therapeutic and/or pharmaceutical compounds (col. 16). Purchio teaches that the chondrocytes can be isolated from articular or costal cartilage (col.11 line 62-65).

Purchio does not teach the method wherein the cartilage is cultured in the manner claimed. However, Masuda teaches methods for culturing an engineered cartilage tissue, wherein isolated chondrocytes are cultured in alginate for a time to allow a cell associated matrix to form (col.4 line 30-35). The cell associated matrix has at least about 5 mg/cc3 aggrecan, a ratio of aggrecan to hyaluronan of 10:1 to 200:1, and a ratio of aggrecan to collagen of about 1:1 to 10: (col.4 line 35-40). The chondrogenic cells are cultured on a semi-permeable membrane in the presence of growth factors (col. 4 line 44-46). The chondrocytes may be isolated from articular cartilage, fibrocartilage, costal, nasal, auricular, tracheal, epiglottic, thyroid, arytenoid or cricoid cartilages (col.5 line 31-38) and the resulting matrix comprises aggrecan, collagn II, IX, XI and hyaluronan (col.6 line 61-64). At the time of the claimed invention, it would have been obvious to one of ordinary skill in the art to culture the cartilage of Purchio using the methods of Masuda because it was a known method for culturing cartilage tissues. Moreover, at the time of the claimed invention, one of ordinary skill in the art would have been motivated by common culture practices in the art to culture the cartilage of Purchio via the methods of Masuda with a reasonable expectation for successfully determining the effects of test agents on cartilage cultures. Although Masuda does not teach the fibrocartilage is from a ligament, tendon, meniscus or intervertebral discs, they were each well known sources of fibrocartilage. Moreover at the time of the claimed invention, it would have been well within the purview of one of

Art Unit: 1651

ordinary skill in the art to use any of the above sources as the source of fibrocartilage in the methods of Masuda.

8. Claims 1 - 12, 14, 17 - 28 and 33 are rejected under 35 U.S.C. 103(a) as being unpatentable over Hunziker (US 5368858 A) in view of Masuda.

Applicant claims a method for determining the effect of a test agent on a tissue engineered cartilage matrix, the method comprising culturing an engineered cartilage tissue (ECT), contacting test agents with cells or tissues of the ECT, and measuring the effect that the agents have on the ECT or cells thereof. The ECT is cultured by culturing isolated chondrogenic cells for a time sufficient to form a chondrogenic cell-associated matrix, and culturing the cells with the cell-associated matrix on a semipermeable membrane in the presence of a growth factor for a time sufficient to form an ECT. The cells or tissues are selected from isolated chondrogenic cells, the chondrogenic cells during culture, the cells and cell-associated matrix before or after culturing, and the ECT. Alternatively, the cells are selected from isolated chondrogenic cells, the chondrogenic cells during culture, the cells and cell-associated matrix before or after culturing, and the ECT in the presence of a known modulator of cartilage tissue. The chondrogenic cell-associated matrix comprises aggrecan, collagen types II, IX, XI, matrix proteins, and hyaluronan and the ECT comprises collagens II, IX, XI, hyaluronan, and at least about 5 mg/cc3 aggrecan, the ratio of aggrecan: hyaluronan is about 10:1-200:1, and the ration of aggrecan: collagen is about 1:1-10:1. The isolated chondrogenic cells are isolated from articular cartilage, costal, nasal, auricular, tracheal, epiglottic, thyroid, arytenoids or cricoid

cartilages, specifically fibrocartilage from ligament, tendon, meniscus, or intervertebral disc.

Art Unit: 1651

The chondrogenic cells are cultured on an alginate medium. The measuring comprises measuring the amount of proteoglycan in the ECT, is performed without addition of extrinsic radioactivity, the ECT is enzymatically degraded and stained with a dye. The method further comprises identifying one or more test agents that have desirable characteristics and producing the agents as a therapeutic drug.

Hunziker teaches a method wherein engineered cartilage is exposed to therapeutic test agents to observe its effects on the cartilage (example 1). The cartilage is fixed and stained for measuring the amount of proteoglycan (example 1).

Hunziker does not teach the method wherein the cartilage is cultured in the manner claimed. However, Masuda teaches methods for culturing an engineered cartilage tissue, wherein isolated chondrocytes are cultured in alginate for a time to allow a cell associated matrix to form (col.4 line 30-35). The cell associated matrix has at least about 5 mg/cc3 aggrecan, a ratio of aggrecan to hyaluronan of 10:1 to 200:1, and a ratio of aggrecan to collagen of about 1:1 to 10: (col.4 line 35-40). The chondrogenic cells are cultured on a semi-permeable membrane in the presence of growth factors (col. 4 line 44-46). The chondrocytes may be isolated from articular cartilage, fibrocartilage, costal, nasal, auricular, tracheal, epiglottic, thyroid, arytenoid or cricoid cartilages (col.5 line 31-38) and the resulting matrix comprises aggrecan, collagn II, IX, XI and hyaluronan (col.6 line 61-64). At the time of the claimed invention, it would have been obvious to one of ordinary skill in the art to culture the cartilage of Hunziker using the methods of Masuda because it was a known method for culturing cartilage tissues. Moreover, at the time of the claimed invention, one of ordinary skill in the art would have been motivated by

common culture practices in the art to culture the cartilage of Hunziker via the methods of

Art Unit: 1651

Masuda with a reasonable expectation for successfully determining the effects of test agents on cartilage cultures. Although Masuda does not teach the fibrocartilage is from a ligament, tendon, meniscus or intervertebral discs, they were each well known sources of fibrocartilage. Moreover at the time of the claimed invention, it would have been well within the purview of one of ordinary skill in the art to use any of the above sources as the source of fibrocartilage in the methods of Masuda.

Claims 1 - 8, 17 - 24, 29 - 30 and 35 are rejected under 35 U.S.C. 103(a) as being 9. unpatentable over Saito (July 1999) in view of Masuda.

Applicant claims a method for determining the effect of a test agent on a tissue engineered cartilage matrix, the method comprising culturing an engineered cartilage tissue (ECT), contacting test agents with cells or tissues of the ECT, and measuring the effect that the agents have on the ECT or cells thereof. The ECT is cultured by culturing isolated chondrogenic cells for a time sufficient to form a chondrogenic cell-associated matrix, and culturing the cells with the cell-associated matrix on a semipermeable membrane in the presence of a growth factor for a time sufficient to form an ECT. The cells or tissues are selected from isolated chondrogenic cells, the chondrogenic cells during culture, the cells and cell-associated matrix before or after culturing, and the ECT. Alternatively, the cells are selected from isolated chondrogenic cells, the chondrogenic cells during culture, the cells and cell-associated matrix before or after culturing, and the ECT in the presence of a known modulator of cartilage tissue. The chondrogenic cell-associated matrix comprises aggrecan, collagen types II, IX, XI, matrix proteins, and hyaluronan and the ECT comprises collagens II, IX, XI, hyaluronan, and at least

Art Unit: 1651

about 5 mg/cc3 aggrecan, the ratio of aggrecan: hyaluronan is about 10:1 – 200:1, and the ration of aggrecan: collagen is about 1:1 – 10:1. The isolated chondrogenic cells are isolated from articular cartilage, costal, nasal, auricular, tracheal, epiglottic, thyroid, arytenoids or cricoid cartilages, specifically fibrocartilage from ligament, tendon, meniscus, or intervertebral disc. The chondrogenic cells are cultured on an alginate medium. The modulator of the ECT tissue is a matrix stimulating agent, cytokine or TNF-alpha wherein the cytokine is interleukin 1 (IL-1). Finally, the culturing and contacting step occur in the same well of a multi well plate.

Saito teaches culturing cartilage in multi well plates in the presences of IL-1 alpha, wherein the effects of the test agent were measured (p.727).

Saito does not teach the method wherein the cartilage is cultured in the manner claimed. However, Masuda teaches methods for culturing an engineered cartilage tissue, wherein isolated chondrocytes are cultured in alginate for a time to allow a cell associated matrix to form (col.4 line 30-35). The cell associated matrix has at least about 5 mg/cc3 aggrecan, a ratio of aggrecan to hyaluronan of 10:1 to 200:1, and a ratio of aggrecan to collagen of about 1:1 to 10: (col.4 line 35-40). The chondrogenic cells are cultured on a semi-permeable membrane in the presence of growth factors (col. 4 line 44-46). The chondrocytes may be isolated from articular cartilage, fibrocartilage, costal, nasal, auricular, tracheal, epiglottic, thyroid, arytenoid or cricoid cartilages (col.5 line 31-38) and the resulting matrix comprises aggrecan, collagn II, IX, XI and hyaluronan (col.6 line 61-64). At the time of the claimed invention, it would have been obvious to one of ordinary skill in the art to culture the cartilage of Saito using the methods of Masuda because it was a known method for culturing cartilage tissues. Moreover, at the time of the claimed

invention, one of ordinary skill in the art would have been motivated by common culture

Art Unit: 1651

practices in the art to culture the cartilage of Saito via the methods of Masuda with a reasonable expectation for successfully determining the effects of test agents on cartilage cultures.

Although Masuda does not teach the fibrocartilage is from a ligament, tendon, meniscus or intervertebral discs, they were each well known sources of fibrocartilage. Moreover at the time of the claimed invention, it would have been well within the purview of one of ordinary skill in the art to use any of the above sources as the source of fibrocartilage in the methods of Masuda.

10. Claims 1 – 11, 17 – 27 and 29 – 30 are rejected under 35 U.S.C. 103(a) as being unpatentable over Huch et al (1997) in view of Masuda.

Applicant claims a method for determining the effect of a test agent on a tissue engineered cartilage matrix, the method comprising culturing an engineered cartilage tissue (ECT), contacting test agents with cells or tissues of the ECT, and measuring the effect that the agents have on the ECT or cells thereof. The ECT is cultured by culturing isolated chondrogenic cells for a time sufficient to form a chondrogenic cell-associated matrix, and culturing the cells with the cell-associated matrix on a semipermeable membrane in the presence of a growth factor for a time sufficient to form an ECT. The cells or tissues are selected from isolated chondrogenic cells, the chondrogenic cells during culture, the cells and cell-associated matrix before or after culturing, and the ECT. Alternatively, the cells are selected from isolated chondrogenic cells, the chondrogenic cells during culture, the cells and cell-associated matrix before or after culturing, and the ECT in the presence of a known modulator of cartilage tissue. The chondrogenic cell-associated matrix comprises aggrecan, collagen types II, IX, XI, matrix proteins, and hyaluronan and the ECT comprises collagens II, IX, XI, hyaluronan, and at least

Art Unit: 1651

about 5 mg/cc3 aggrecan, the ratio of aggrecan: hyaluronan is about 10:1 – 200:1, and the ration of aggrecan: collagen is about 1:1 – 10:1. The isolated chondrogenic cells are isolated from articular cartilage, costal, nasal, auricular, tracheal, epiglottic, thyroid, arytenoids or cricoid cartilages, specifically fibrocartilage from ligament, tendon, meniscus, or intervertebral disc. The chondrogenic cells are cultured on an alginate medium, the measuring comprises measuring the amount of proteoglycan in the ECT, enzymatically degrading the ECT and is performed without addition of extrinsic radioactivity. The method further comprises identifying one or more test agents that have desirable characteristics and producing the agents as a therapeutic drug. The culturing of ECT and the contacting the cells with the test agent occurs in the same well of a multi-well plate. The modulator of the ECT tissue is a matrix stimulating agent, cytokine or TNF-alpha wherein the cytokine is interleukin 1.

Huch teaches methods for culturing articular chondrocytes in an alginate medium in the presence of a test agent, IL-1, wherein proteoglycan was measured (abstract). Specifically, the cartilage was degraded with enzymes, the chondrocytes were cultured with alginate in a multiwell plate in the presence of IL-1, and the amount of proteoglycan was measured (p. 2158).

Huch does not teach the method wherein the cartilage is cultured in the manner claimed. However, Masuda teaches methods for culturing an engineered cartilage tissue, wherein isolated chondrocytes are cultured in alginate for a time to allow a cell associated matrix to form (col.4 line 30-35). The cell associated matrix has at least about 5 mg/cc3 aggrecan, a ratio of aggrecan to hyaluronan of 10:1 to 200:1, and a ratio of aggrecan to collagen of about 1:1 to 10: (col.4 line 35-40). The chondrogenic cells are cultured on a semi-permeable membrane in the presence of growth factors (col. 4 line 44-46). The chondrocytes may be isolated from articular cartilage,

Art Unit: 1651

fibrocartilage, costal, nasal, auricular, tracheal, epiglottic, thyroid, arytenoid or cricoid cartilages (col.5 line 31-38) and the resulting matrix comprises aggrecan, collagn II, IX, XI and hyaluronan (col.6 line 61-64). At the time of the claimed invention, it would have been obvious to one of ordinary skill in the art to culture the cartilage of Huch using the methods of Masuda because it was a known method for culturing cartilage tissues. Moreover, at the time of the claimed invention, one of ordinary skill in the art would have been motivated by common culture practices in the art to culture the cartilage of Huch via the methods of Masuda with a reasonable expectation for successfully determining the effects of test agents on cartilage cultures.

Although Masuda does not teach the fibrocartilage is from a ligament, tendon, meniscus or intervertebral discs, they were each well known sources of fibrocartilage. Moreover at the time of the claimed invention, it would have been well within the purview of one of ordinary skill in the art to use any of the above sources as the source of fibrocartilage in the methods of Masuda.

11. Claims 1 - 8, 10, 14, 17 - 24, 26 and 33 are rejected under 35 U.S.C. 103(a) as being unpatentable over Lansbury et al. (WO 94/28889) in view of Masuda.

Applicant claims a method for determining the effect of a test agent on a tissue engineered cartilage matrix, the method comprising culturing an engineered cartilage tissue (ECT), contacting test agents with cells or tissues of the ECT, and measuring the effect that the agents have on the ECT or cells thereof. The ECT is cultured by culturing isolated chondrogenic cells for a time sufficient to form a chondrogenic cell-associated matrix, and culturing the cells with the cell-associated matrix on a semipermeable membrane in the presence of a growth factor

Art Unit: 1651

chondrogenic cells, the chondrogenic cells during culture, the cells and cell-associated matrix before or after culturing, and the ECT. Alternatively, the cells are selected from isolated chondrogenic cells, the chondrogenic cells during culture, the cells and cell-associated matrix before or after culturing, and the ECT in the presence of a known modulator of cartilage tissue. The chondrogenic cell-associated matrix comprises aggrecan, collagen types II, IX, XI, matrix proteins, and hyaluronan and the ECT comprises collagens II, IX, XI, hyaluronan, and at least about 5 mg/cc3 aggrecan, the ratio of aggrecan: hyaluronan is about 10:1 – 200:1, and the ration of aggrecan: collagen is about 1:1 – 10:1. The isolated chondrogenic cells are isolated from articular cartilage, costal, nasal, auricular, tracheal, epiglottic, thyroid, arytenoids or cricoid cartilages, specifically fibrocartilage from ligament, tendon, meniscus, or intervertebral disc. The chondrogenic cells are cultured on an alginate medium, the measuring is performed without addition of extrinsic radioactivity and the method further comprises identifying one or more test agents that have desirable characteristics and producing the agents as a therapeutic drug.

Lansbury teaches methods for screening the effects of agents on cartilage cultures wherein a chondrocyte cell culture is incubated (or contacted) with the test agent and the effects are measured (claim 34). The method is used to identify agents with desirable, therapeutic characteristics, specifically the ability to repair damaged cartilage (claim 34).

Lansbury does not teach the method wherein the cartilage is cultured in the manner claimed. However, Masuda teaches methods for culturing an engineered cartilage tissue, wherein isolated chondrocytes are cultured in alginate for a time to allow a cell associated matrix to form (col.4 line 30-35). The cell associated matrix has at least about 5 mg/cc3 aggrecan, a ratio of aggrecan to hyaluronan of 10:1 to 200:1, and a ratio of aggrecan to collagen of about 1:1

Art Unit: 1651

to 10: (col.4 line 35-40). The chondrogenic cells are cultured on a semi-permeable membrane in the presence of growth factors (col. 4 line 44-46). The chondrocytes may be isolated from articular cartilage, fibrocartilage, costal, nasal, auricular, tracheal, epiglottic, thyroid, arytenoid or cricoid cartilages (col.5 line 31-38) and the resulting matrix comprises aggrecan, collagn II, IX, XI and hyaluronan (col.6 line 61-64). At the time of the claimed invention, it would have been obvious to one of ordinary skill in the art to culture the cartilage of Lansbury using the methods of Masuda because it was a known method for culturing cartilage tissues. Moreover, at the time of the claimed invention, one of ordinary skill in the art would have been motivated by common culture practices in the art to culture the cartilage of Lansbury via the methods of Masuda with a reasonable expectation for successfully determining the effects of test agents on cartilage cultures. Although Masuda does not teach the fibrocartilage is from a ligament, tendon, meniscus or intervertebral discs, they were each well known sources of fibrocartilage. Moreover at the time of the claimed invention, it would have been well within the purview of one of ordinary skill in the art to use any of the above sources as the source of fibrocartilage in the methods of Masuda.

Any inquiry concerning this communication or earlier communications from the examiner should be directed to Ruth A. Davis whose telephone number is 703-308-6310. The examiner can normally be reached on M-H (7:00-4:30); altn. F (7:00-3:30).

If attempts to reach the examiner by telephone are unsuccessful, the examiner's supervisor, Michael Wityshyn can be reached on 703-308-0196. The fax phone numbers for the

Page 18

Application/Control Number: 10/054,710

Art Unit: 1651

organization where this application or proceeding is assigned are 703-308-4242 for regular communications and 703-308-4242 for After Final communications.

Any inquiry of a general nature or relating to the status of this application or proceeding should be directed to the receptionist whose telephone number is 703-308-0196.

Ruth A. Davis; rad May 1, 2003

> LEÓN/B./LANKFORD, JR. PRIMARY EXAMINER